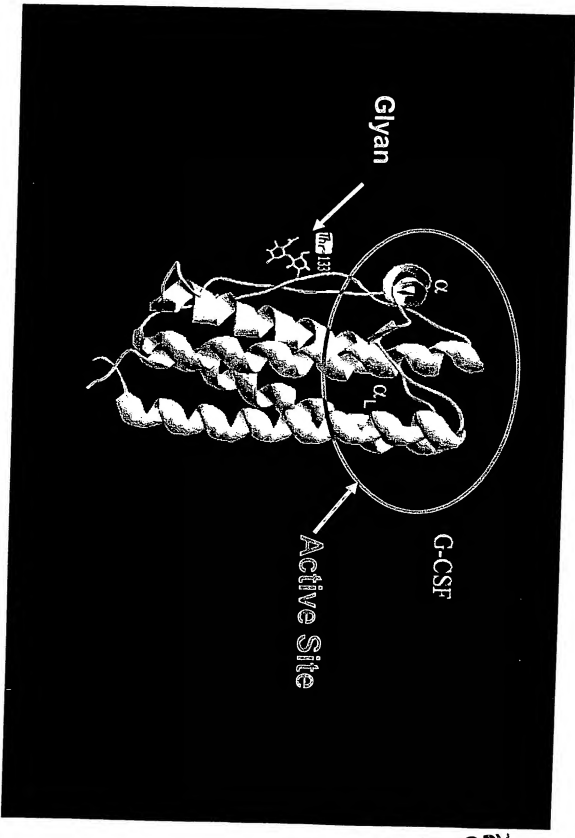
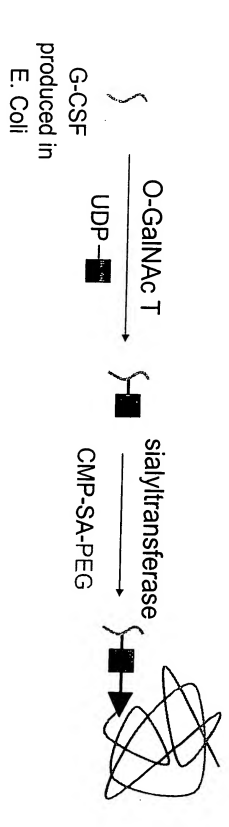
FIGURE 1



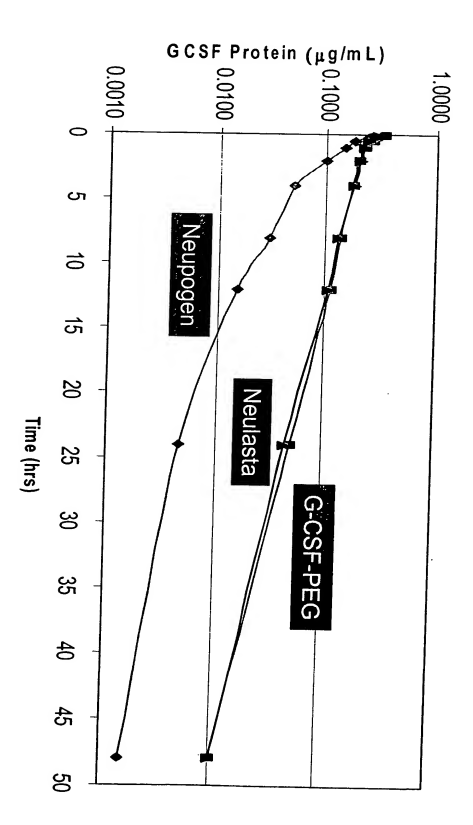
BEST AVAILABLE COPY

FIGURE 2



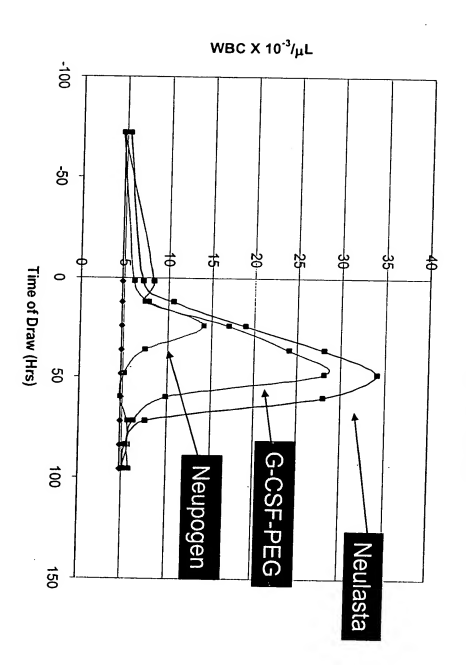
PEST AVAILABLE COPY

FIGURE 3



IV Bolus injection of [125]]-labeled proteins in rats

FIGURE 4



# Mouse WBC Response to GCSF Variants (250 μg/Kg) Administered i.v. at 0 hours

### 175 amino acid variant

MTPLGPASSLPQSFLLKCLEQVRKIQGDGAALQEKLCA
TYKLCHPEELVLLGHSLGIPWAPLSSCPSQALQLAGCLS
QLHSGLFLYQGLLQALEGISPELGPTLDTLQLDVADFAT
TIWQQMEELGMAPALQPTQGAMPAFASAFQRRAGGVL
VASHLQSFLEVSYRVLRHLAQP (SEQ ID NO: 1).

### 174 amino acid variant

TPLGPASSLPQSFLLKCLEQVRKIQGDGAALQEKLCATY KLCHPEELVLLGHSLGIPWAPLSSCPSQALQLAGCLSQL HSGLFLYQGLLQALEGISPELGPTLDTLQLDVADFATTI WQQMEELGMAPALQPTQGAMPAFASAFQRRAGGVLV ASHLQSFLEVSYRVLRHLAQP (SEQ ID NO: 2).

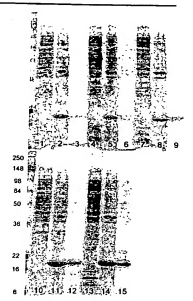
FIGURE 7

# Testing growth conditions - washing IBs

	37°C,	1mM (final) IPT	G	IB yields (~0.7-1g/L)
L	Media	Buffered pH	Time	wet IB pellet (mg)
1.	LB	n/a	5hr	120
2.	TB	7.4 (K-PO <sub>4</sub> )	5hr	230
3.	ТВ	6 (MES)	5hr	170
4.	LB	n/a	o/n	190
5.	тв	7.4 (K-PO <sub>4</sub> )	o/n	250

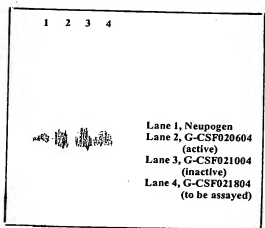
Cell samples, loaded equally by ug (better expression of overnights more apparent)

- #1 lysate, supe
- 10. #4 lysate, supe
- #1 lysate, pellet #1 final IB
- 11. #4 lysate, pellet 12. #4 final IB
- 13. #5 lysate, supe
- #2 lysate, supe #2 lysate, pellet
- 14. #5 lysate, pellet
- #2 final IB #3 lysate, supe
- 15. #5 final IB
- #3 lysate, pellet #3 final IB



Purified IB's are shown in lanes 3, 6, 9, 12 and 15. Note IB lanes loaded at approximately ½ the amount of the preceding lane (based on predicted pellet weight).

# Western Blot Analysis of Refolded G-CSF Native Polyacrylamide Gel Electrophoresis



G-CSF020604 and G-CSF021004 came from the same batch of refolding reaction. The only difference was that G-CSF021004 was saved from the flowthough of G-CSF020604 by adjusting pH and reloading onto SP Sepharose. G-CSF021804 was a separate refolding batch.

After SP-sepharose you see only one band by Coomassie stain or Western.

# FIGURE 11A

Protein	Organism	EC#	GenBai	nk / GenPept	SwissPro	
At1g08280	Arabidopsis thaliana	n.d.	AC011438	AAF18241.1	Q84W00	// 3
	, addicopolo alamana	, Ia.	BT004583			
		Ì		0 NP_172305.		
At1g08660/F22O13.14	Arabidopsis thaliana	n.d.		AAF99778.1		
geecco 220 (0	i i doldopsis tranaria	, j.i.d.	AY064135			
	<u>.</u>	1		AAM70516.1	QSFRRS	
				0 NP_172342.		
			NM 18060	9 NP_850940.	:1	
At3g48820/T21J18 90	Arabidopsis thaliana	n.d.	AV080580	AAL85966.1	Q8RY00	
	, washaopala wanana	11.0.	AY133816			
			AL132963			
				1NP_190451.		
α-2,3-sialyltransferase	Bos taurus	n.d.	AJ584673	CAE48298.1		
(ST3GAL-IV)	200 144.40	Ju.G.	73304073	CAL40250.1		
cc-2,3-sialyltransferase	Bos taurus	n.d.	AJ585768	CAE51392.1	+	
(St3Gal-V)	Dos tauras	ii.u.	A3303700	CAE51392.1	İ	
α-2,6-sialyltransferase	Bos taurus	n.d.	AJ620651	CAF05850.1	<del> </del>	
(Siat7b)	Dos tautus	11.u.	A3020651	CAF03630.1		
α-2,8-sialyltransferase	Bos taurus	2 4 00 0	AJ699418	CAC27990 4	<del>-</del>	
(SIAT8A)	DOS taurus	2.4.99.0	MJ099418	CAG27880.1		
α-2,8-sialyltransferase	Bos taurus	n d	A 1600404	CACOZOGO	<del> </del>	
(Siat8D)	BUS laurus	n.d.	AJ699421	CAG27883.1	1	
α-2,8-sialyltransferase	Bos taurus	-	A 170 1505	0400000	<del> </del>	
ST8Siα-III (Siat8C)	pos laurus	n.d.	AJ704563	CAG28696.1		
CMP α-2,6-	Pos tours	0.4.00	146111	<del> </del>		
sialyltransferase	Bos taurus	2.4.99.1	Y15111	CAA75385.1	O18974	
ST6Gal I)			NM_177517	NP_803483.1	1	
					<u> </u>	
sialyltransferase 8	Bos taurus	n.d.	AF450088	AAL47018.1	Q8WN13	
fragment)						
sialyltransferase	Bos taurus	n.d.	AJ748841	CAG44450.1		
ST3Gal-II (Siat4B)			L			
sialyltransferase	Bos taurus	n.d.	AJ748842	CAG44451.1		
ST3Gal-III (Siat6)						
sialyltransferase	Bos taurus	n.d.	AJ748843	CAG44452.1		
ST3Gal-VI (Siat10)						
ST3Gal I	Bos taurus	n.d.	AJ305086	CAC24698.1	Q9BEG4	
St6GalNAc-VI	Bos taurus	n.d.	AJ620949	CAF06586.1		
CDS4	Branchiostoma	n.d.	AF391289	AAM18873.1	Q8T771	
	floridae					
oolysialyltransferase	Cercopithecus	2.4.99	AF210729	AAF17105.1	Q9TT09	
PST) (fragment)	aethiops					
T8Sia IV			ĺ			
oolysialyltransferase	Cercopithecus	2.4.99	AF210318	AAF17104.1	Q9TT10	
STX) (fragment)	aethiops					
T8Sia II						
r-2,3-sialyltransferase	Ciona intestinalis	n.d.	AJ626815	CAF25173.1		
T3Gal I (Siat4)						
c-2,3-sialyltransferase	Ciona savignyi	n.d.	AJ626814	CAF25172.1		
T3Gal I (Siat4)			5020017			
c-2,8-	Cricetulus griseus	2.4.99		AAE28634	Q64690	
olysialyltransferase				CAA86822.1	W0+030	
T8Sia IV			70001	U/V\00022.1		
Sal β-1,3/4-GIcNAc α-	Cricetulus griseus	n.d.	AY266675	AAP22942.1	Q80WL0	
3-sialyltransferase	2Joitalad gilodad	····u.	A1200073		⊶00VVLU	
3Gal I		ļ	ŀ	j		
Sal β.1,3/4-GlcNAc α-	Cricetulus griseus		AV/000070	4 4 DOOG 12 1		
2 cialultennesses	Circeiuius griseus	n.d.	AY266676	AAP22943.1	280WK9	
3-sialyltransferase	1	ŀ	ļ			İ
3Gal II (fragment)						
-2,3-sialyltransferase	Danio rerio r	n.d.	AJ783740	CAH04017.1		
「3Gal I (Siat4)						
-2,3-sialyltransferase [3Gal II (Siat5)	Danio rerio r	ı.d. /	AJ783741 (	CAH04018.1		$\neg$
(161-111 (61.46)						

### FIGURE 11B

Protein	Organism	EC#	GenBa	nk / GenPept	SwissProt	
α-2,3-sialyltransferase ST3Gal III (Siat6)	Danio rerio	n.d.	AJ626821	CAF25179.1		/ 31
α-2,3-sialyltransferase ST3Gal IV (Siat4c)	Danio rerio	n.d.	AJ744809	CAG32845.1	1	
α-2,3-sialyltransferase ST3Gal V-r (Siat5- related)	Danio rerio	n.d.	AJ783742	CAH04019.1	1	
α-2,6-sialyltransferase ST6Gal I (Siat1)	Danio rerio	n.d.	AJ744801	CAG32837.1	1	
α-2,6-sialyltransferase  ST6GalNAc II (Siat7B)	Danio rerio	n.d.	AJ634459	CAG25680.1		
α-2,6-sialyltransferase ST6GalNAc V (Siat7E) (fragment)	Danio rerio	n.d.	AJ646874	CAG26703.1		
α-2,6-sialyltransferase ST6GalNAc VI (Siat7F) (fragment)	Danio rerio	n.d.	AJ646883	CAG26712.1		
α-2,8-sialyltransferase ST8Sia I (Siat 8A) (fragment)	Danio rerio	n.d.	AJ715535	CAG29374.1		<u> </u>
α-2,8-sialyltransferase ST8Sia III (Siat 8C) (fragment)	Danio rerio	n.d.	AJ715543	CAG29382.1		
	Danio rerio	n.d.	AJ715545	CAG29384.1		
α-2,8-sialyltransferase ST8Sia V (Siat 8E) (fragment)	Danio rerio	n.d.	AJ715546	CAG29385.1		-
α-2,8-sialyltransferase ST8Sia VI (Siat 8F) (fragment)	Danio rerio	n.d.	AJ715551	CAG29390.1		
Galactosamide α-2,6- sialyltransferase II (ST6Gal II)	Danio rerio	n.d.	AJ627627	CAF29495.1		
N-glycan α-2,8- sialyltransferase	Danio rerio	n.d.	BC050483 AY055462 NM 153663		Q7ZU51 Q8QH83	
ST3Gal III-related siat6r)	Danio rerio	n.d.	BC053179 AJ626820		Q7T3B9	
St3Gal-V	Danio rerio	n.d.	AJ619960	CAF04061.1	1	
st6GaINAc-VI	Danio rerio	n.d.	BC060932 AJ620947	AAH60932.1 CAF06584.1		
x-2,6-sialyltransferase CG4871) ST6Gal I	Drosophila melanogaster		AE003465 AF218237 AF397532 AE003465 NM_079129	AAF47256.1	Q9GU23 Q9W121	-
x-2,3-sialyltransferase ST3Gal-VI)	Gallus gallus	n.d.	AJ585767	CAE51391.1 CAF25503.1		
x-2,3-sialyltransferase T3Gal I	Gallus gallus	2.4.99.4	X80503		Q11200	
c-2,3-sialyltransferase T3Gal IV (fragment)	Gallus gallus				O73724	$\dashv$
c-2,3-sialytransferase ST3GAL-II)	Gallus gallus	n.d.	AJ585761	CAE51385.2		-
c-2,6-sialyltransferase Siat7b)	Gallus gallus	n.d.	AJ620653	CAF05852.1		$\dashv$
r-2,6-sialyltransferase T6Gal I	Gallus gallus	2.4.99.1		CAA53235.1 NP_990572.1	92182	
-2,6-sialyltransferase	Gallus gallus	2.4.99.3			92183	$\dashv$

### FIGURE 11C

Protein	Organism	EC#	GenBan	k / GenPept	SwissProt	PDB / 3D
ST6GalNAc I			-	AAE68029.1	<del>                                     </del>	1 30
	ļ		X74946	CAA52902.1		
		i	NM_205240	NP_990571.1		
α-2,6-sialyltransferase	Gallus gallus	2.4.99		AAE68030.1	Q92184	
ST6GalNAc II			NM_205233	CAA54813.1		
				NP_990564.1		
α-2,6-sialyltransferase	Gallus gallus	n.d.	AJ634455	CAG25677.1		
ST6GalNAc III (SIAT7C)					i	
(fragment)						
α-2,6-sialyltransferase	Gallus gallus	n.d.	AJ646877	CAG26706.1		
ST6GalNAc V (SIAT7E) (fragment)					1	
α-2,8-sialyltransferase	Co#		-			
(GD3 Synthase) ST8Sia	Gallus gallus	2.4.99	U73176	AAC28888.1	P79783	
(GD3 Synthase) S16Sia					ļ	
α-2,8-sialyltransferase	Callus callus		1000440	040070044	<del> </del>	
(SIAT8B)	Gallus gallus	n.d.	AJ699419	CAG27881.1	ļ	
α-2,8-sialyltransferase	College college		4 1000 400	0.000000	ļ	
(SIAT8C)	Gallus gallus	n.d.	AJ699420	CAG27882.1		
α-2,8-sialyltransferase	Gallus gallus	n.d.	AJ699424	CA C07000 4		
(SIAT8F)	Gallus gallus	n.a.	AJ099424	CAG27886.1	l	
α-2,8-syalyltransferase	Gallus gallus	n.d.	AJ704564	CAG28697.1		
ST8Sia-V (SIAT8C)	Ganus ganus	n.a.	AJ704364	CAG28697.1	1	- 1
galactosamide α-2,6-	Gallus gallus	n.d.	AJ627629	CAF29497.1		
sialyltransferase II	Canas ganas	11.0.	MJ02/029	CAF29497.1		
ST6Gal II)					ļ	
GM3 synthase (SIAT9)	Callerande	0.4.00.0	4)/545055	4.400010.1		
polysialyltransferase	Gallus gallus			AAS83519.1	2 12 2 2 2	
ST8Sia IV	Gallus gallus	2.4.99	AF008194	AAB95120.1	O42399	- 1
x-2,3-sialyltransferase	Homo sapiens	0.4.00.4	1.00555	44400404		
ST3Gal I	Homo Sapiens	2.4.99.4			Q11201	1
, , , , , , , , , , , , , , , , , , ,	İ				O60677	
ł	i i			AAC37574.1   AAD39238.1	Q9UN51	
ļ				AAG29876.1		
				AAH18357.1		
ĺ	i			NP_003024.1		j
			NM_173344	NP 775479 1		1
x-2,3-sialyltransferase	Homo sapiens	2.4.99.4			Q16842	-
ST3Gal II	·				O00654	
	İ			CAA65447.1		
			NM_006927	NP 008858.1		
x-2,3-sialyltransferase	Homo sapiens	2.4.99.6			Q11203	
T3Gal III (SiaT6)			BC050380		Q86UR6	- [
	İ			AAO13859.1	Q86UR7	1
<b>†</b>	İ		AF425852 /		Q86UR8	- 1
ł			AF425853	AO13861.1	286UR9	- 1
		, ,	AF425854	AO13862.1	286US0	
ļ					286US1	
					286US2	
					28IX43	
	i				28IX44	
	ĺ				28IX45	
	ľ	1		AO13868.1		
j				AO13869.1 C	281X47	
	ĺ					
1	1		AF425864 A	AO13871.1 C AO13872.1 C		
Į.	1				281X50 281X51	
ļ	1			AO13873.1 C		
					181X53	
	1			AO38806.1 Q		
	1				181X55	
	· ·				81X56	ļ
		<u> </u>				

### FIGURE 11D

Protein	Organism		Composite / Composite	lo
Tiotem	Organism	EC#	GenBank / GenPept	SwissProt PDE / 3D
			AY167995 AAO38809.1 AY167996 AAO38810.1 AY167997 AAO38811.1 AY167998 AAO38812.1 NM_006279 NP_006270. NM_174964 NP_777624.1 NM_174966 NP_777625.1 NM_174967 NP_777627.1 NM_174969 NP_777630.1 NM_174970 NP_777632.1	Q8IX58
α-2,3-sialyltransferase ST3Gal IV	Homo sapiens	2.4.99	L23767 AF035249 BC010645 AY040826 AF516602 AF516603 AF516604 AF525084 X74570 CR456858 NM_006278 NP_006269.1	Q11206 O60497 Q96QQ9 Q8N6A6 Q8N6A7 Q8NFD3 Q8NFG7
α-2,3-sialyltransferase ST3Gal VI	Homo sapiens	2.4.99.4	AF119391 AAD39131.1 BC023312 AAH23312.1 AB022918 BAA77609.1 AX877828 CAE89895.1 AX886023 CAF00161.1 NM_006100 NP_006091.1	Q9Y274
α-2,6-sialyltransferase (ST6Gal II ; KIAA1877)	Homo sapiens	n.d.	BC008680 AAH08680.1 AB058780 BAB47506.1 AB059555 BAC24793.1	Q86Y44 Q8IUG7 Q96HE4 Q96JF0
	Homo sapiens	n.d.	BC059363 AAH59363.1	Q8N259 Q8NDV1
α-2,6-sialyltransferase (ST6GalNAc V)	Homo sapiens	n.d.	BC001201 AAH01201.1 AK056241 BAB71127.1 CAB72344.1 CAD45372.1 NM 030965 NP 112227.1	Q9BVH7
α-2,6-sialyltransferase (SThM) ST6GalNAc II	Homo sapiens			Q9UJ37 Q12971
α-2,6-sialyltransferase ST6Gal I	Homo sapiens	2.4.99.1		215907
∝-2,6-sialyltransferase ST6GalNAc I	Homo sapiens	2.4.99.3 E	BC022462 AAH22462.1 C AY096001 AAM22800.1 C	28TBJ6 29NSC7 29NXQ7

### FIGURE 11E

De .	FIGUR		· · · · · · · · · · · · · · · · · · ·		
Protein	Organism	EC#	GenBank / GenPept	SwissProt PD	
			NM_018414 NP_060884		
α-2,8-	Homo sapiens	2.4.99	L41680 AAC41775.1		
polysialyltransferase			BC027866 AAH27866.1		
ST8Sia IV	1		BC053657 AAH53657.	Q92693	
			NM_005668 NP_005659.	1	
α-2,8-sialyltransferase	Homo sapiens	2.4.99.8	L32867 AAA62366.1		
(GD3 synthase) ST8Sia			L43494 AAC37586.1		
	ĺ		BC046158 AAH46158.1	Q93064	
		1	- AAQ53140.1		
			AY569975 AAS75783.1		
			D26360 BAA05391.1		
	ļ		X77922 CAA54891.1		
er O O cial than a f			NM_003034 NP_003025.		
α-2,8-sialyltransferase	Homo sapiens	2.4.99	L29556 AAA36613.1		
ST8Sia II			U82762 AAB51242.1		
			U33551 AAC24458.1		
			BC069584 AAH69584.1		
er 2 8 sighttenneformen			NM_006011 NP_006002.		
α-2,8-sialyltransferase ST8Sia III	Homo sapiens	2.4.99	AF004668 AAB87642.1		
5100ia III			AF003092 AAC15901.2		
α-2,8-sialyltransferase	Home as	0.4.00	NM_015879 NP_056963.		
ST8Sia V	Homo sapiens	2.4.99	U91641 AAC51727.1		
J. Jola v			CR457037 CAG33318.1		
ENSP00000020221		_	NM_013305 NP_037437.1		
fragment)		n.d.	AC023295 -		
actosylceramide α-2,3-	Homo sapiens	2 4 90 0	AF105026 AAD14634.1	COLINIDA	
ialyltransferase	i iome dapiens		AF119415 AAF66146.1	Q9UNP4 O94902	
ST3Gal V)			BC065936 AAH65936.1	094902	
,	}		AY152815 AAO16866.1		
	1		AAP65066 AAP65066.1		
	Ì		AY359105 AAQ89463.1		
İ	1		AB018356 BAA33950.1		
	ļ		AX876536 CAE89320.1		
			NM_003896 NP_003887.2		
V-	Homo sapiens	2.4.99	BC006564 AAH06564.1	Q969X2	
cetylgalactosaminide			BC007802 AAH07802.1	Q9H8A2	
-2,6-sialyltransferase	}		BC016299 AAH16299.1	Q9ULB8	
ST6GalNAc VI)			AY358672 AAQ89035.1		
		1 1	AB035173 BAA87035.1	i	
J	1		AK023900 BAB14715.1		
	[		AJ507293 CAD45373.1		
1	ĺ		X880950 CAE91145.1		
1			CR457318  CAG33599.1	ļ	
			NM_013443 NP_038471.2		
-	Homo sapiens	2.4.99	AF127142 AAF00102.1	Q9H4F1	
cetylgalactosaminide		8		Q9NWU6	
2,6-sialyltransferase		-	AAP63349.1	Q9UKU1	
(ST6GaINAc IV)			B035172 BAA87034.1	Q9ULB9	
.			K000600 BAA91281.1	Q9Y3G3	
ł			'17461   CAB44354.1	Q9Y3G4	
	1		J271734 CAC07404.1		
1	ł	1 5	X061620 CAC24981.1		
			X068265 CAC27250.1		
ļ	ļ		X969252 CAF14360.1		
	İ		M_014403 NP_055218.3	Ì	
TOCIA VII (from the second			M_175039 NP_778204.1		
T8SIA-VI (fragment)	Homo sapiens		J621583 CAF21722.1		
			M_291725 XP_291725.2		
nnamed protein	Homo sapiens	n.d. A	K021929 BAB13940.1	Q9HAA9	
oduct		A	X881696   CAE91353.1		
al β.1,3/4-GlcNAc α-	Mesocricetus	2.4.99.6 A		29QXF6	

# FIGURE 11F

Protein		Organism	EC	# GenBan	k / GenPept	SwissPro	t Pi
2,3-sialyltransferase (ST3Gal III)		auratus			T		1
Gal β 1,3/4-GlcNAc α 2,3-sialyltransferase (ST3Gal IV)	-	Mesocricetus auratus	2.4.99	0.6 AJ245700	CAB53395.	1 Q9QXF5	
GD3 synthase (fragment) ST8Sia I		Mesocricetus auratus	n.d.	AF141657	AAD33879.	1 Q9WUL1	
polysialyltransferase (ST8Sia IV)		Mesocricetus auratus	2.4.99	AJ245701	CAB53396.1	I Q9QXF4	
α-2,3-sialyltransferase ST3Gal I	St3gal1	Mus musculus	2.4.99	AK031344 AK078469	AAF60973.1 BAC27356.1 BAC37290.1	<b>Q11202</b> Q9JL30	
α-2,3-sialyltransferase ST3Gal II	St3gal2	Mus musculus	2.4.99.	NM_009177 4 BC015264 BC066064 AK034554 AK034863 AK053827	AAH15264.1 AAH66064.1 BAC28752.1 BAC28859.1 BAC35543.1 CAA54294.1	1 Q11204 Q8BPL0 Q8BSA0 Q8BSE9 Q91WH6	
α-2,3-sialyltransferase ST3Gal III	St3gal3	Mus musculus	2.4.99	NM_178048   BC006710   AK005053   AK013016   X84234   C	NP_835149.1 AAH06710.1 BAB23779.1 BAB28598.1 CAA59013.1	P97325 Q922X5 Q9CZ48	
α-2,3-sialyltransferase ST3Gal IV	St3gal4	Mus musculus	2.4.99.4	BC050773 A D28941 B AK008543 B AB061305 B X95809 C	IP_033202.2 AH11121.1 AH50773.1 AA06068.1 AB25732.1 AB47508.1 AA65076.1	P97354	
ST3Gal VI	St3gal6	Mus musculus		BC052338 A AB063326 B AK033562 B AK041173 BA	AD39130.1 AH52338.1 AB79494.1 AC28360.1 AC30851.1	Q80UR7 Q8BLV1 Q8VIB3 Q9WVG2	
TOGAINAC II		2 Mus musculus	2.4.99	AK004613 BA X93999 CA X94000 CA	77963 FAH10208.1 CAB00637.1 CAB23410.1 CAB3821.1 CAB3822.1	P70277 Q9DC24 Q9JJM5	
F6Gal I	St6gal1	Mus musculus	2.4.99.1  -       	BC027833  AA  D16106  BA  NK034768  BA  NK084124  BA	E68031.1 Q H27833.1 Q A03680.1 Q C28828.1 C39120.1	8BM62 8K1L1	
6Gal II	t6gal2	Mus musculus	n.d.	B095093 <b>BA</b> 0 K129462 BA0	C38534.1 Q8 C87752.1 C98272.1	BBUU4	
oGainac I		Mus musculus	[2.4.99.3  Y		72137.1 QS	QZ39	+
2,6-sialyltransferase St 6GalNAc III	6galnac3	Mus musculus	n.d. B	K034804 BAC	58387.1 Q9	JJJP5 IWUV2 JHP5	

### FIGURE 11G

Protein		Organism	EC#	GenBar	nk / GenPept	SwissPro	t PDB
				NM 01137	2NP_035502		1,30
α-2,6-sialyltransferase	St6galna	c4 Mus musculus	2.4.99.		AAH56451.1		
ST6GalNAc IV	1			AK085730	BAC39523.1		
			ľ	AJ007310	CAA07446.1		
				Y15779	CAB43507.1		
	İ			Y15780	CAB43514.1	I Q9JHP0	
1				Y19055	CAB93946.1		
İ	1	İ	ĺ	Y19057	CAB93948.1		
er 2.9 sightlespetares	C400-in4				3 NP_035503.		
α-2,8-sialyltransferase (GD3 synthase) ST8Si		Mus musculus	2.4.99.	-	AAA91869.1		
i	a	ļ	ļ	BC024821	AAH24821.1		
•	ļ			AK046188	BAC32625.1		
	ļ		İ	AK052444	BAC34994.1		
			1	X84235	CAA59014.1		
			1	AJ401102	CAC20706.1		
α-2,8-sialyltransferase	St8sia6	Mus musculus	n.d.		NP_035504.		
(ST8Sia VI)	0.03,40	inas mascalas	m.a.	AB059554	BAC01265.1		
(=======				AK085105	BAC39367.1 NP_665837.	Q8K4T1	1
α-2,8-sialyltransferase	St8sia2	Mus musculus	2.4.99	X83562			
ST8Sia II	0.00.02	indo mascaras	2.4.99	X99646	CAA58548.1 CAA67965.1	O35696	l
	Í			X99647	CAA67965.1		
	ĺ	ļ	1	X99648	CAA67965.1		
				X99649	CAA67965.1		Į
	ŀ			X99650	CAA67965.1		l
	1		]	X99651	CAA67965.1	İ	
					NP_033207.1		- 1
α-2,8-sialyltransferase	St8sia4	Mus musculus	2.4.99.8	BC060112	AAH60112.1	Q64692	
ST8Sia IV	1		ļ	AK003690	BAB22941.1	Q8BY70	j
	ļ		ļ	AK041723	BAC31044.1		
	}			AJ223956	CAA11685.1		
		1		X86000	CAA59992.1		
	1	İ	1	Y09484	CAA70692.1		
x-2,8-sialyltransferase	St8sia5	144			NP_033209.1		
ST8Sia V	Siosias	Mus musculus	2.4.99		AAH34855.1	P70126	1
orocia v	1				BAC37354.1	P70127	1
					CAA66642.1	P70128	- 1
	İ		· ·		CAA66643.1	Q8BJW0	
		1	1		CAA66644.1 NP_038694.1	Q8JZQ3	
	ļ	1	1		NP_694764.1		.
			ĺ	NM 177416	NP_803135.1		- 1
x-2,8-sialytransferase	St8sia3	Mus musculus	2.4.99	BC075645		Q64689	
T8Sia III	İ					Q9CUJ6	
		ł		1 1	CAA56665.1	Q3C030	j
					NP_033208.1		
GD1 synthase	St6galnac5	Mus musculus	n.d.	BC055737		Q8CAM7	
ST6GalNAc V)						Q8CBX1	- 1
			1	AB028840		Q9QYJ1	- 1
		}	1	AK034387		Q9R0K6	
		1		AK038434	BAC29997.1		
		1		AK042683	BAC31331.1		
2442 overthere a (= 0.0	010 15			NM_012028	NP_036158.2		j
SM3 synthase (α-2,3- alyltransferase)	St3gal5	Mus musculus	2.4.99.9			088829	
ragal V						Q9CZ65	
. ocal v						Q9QWF9	]
					BAA76467.1		
ł	i	4			BAB28571.1		- 1
					CAA75235.1		
_	St6galnac6	Mus musculus		NM_011375 N	MU20005.1	200000	
etylgalactosaminide	094//1000	mus musculus		BC036985 A		28CDC3	
2,6-sialyltransferase	ľ					28JZW3	j
T6GalNAc VI)						29JM95	-
			ı <i>V</i>	717UJUU40 IB	AC27064.1 C	29R0G9	- 1

### FIGURE 11H

Protein	Organism	EC#	GenBai	nk / GenPept	SwissProt	PD / 31
			NM 01697	3NP_058669.1	<del> </del>	1, 5
M138L	Myxoma virus	n.d.	U46578 AF170726	AAD00069.1		
α-2,3-sialyltransferase (St3Gal-I)	Oncorhynchus mykiss	n.d.	AJ585760	CAE51384.1		
α-2,6-sialyltransferase (Siat1)	Oncorhynchus mykiss	n.d.	AJ620649	CAF05848.1		
α-2,8- polysialyltransferase IV (ST8Sia IV)	Oncorhynchus mykiss	n.d.	AB094402	BAC77411.1	Q7T2X5	
GalNAc α-2,6- sialyltransferase (RtST6GalNAc)	Oncorhynchus mykiss	n.d.	AB097943	BAC77520.1	Q7T2X4	
α-2,3-sialyltransferase ST3Gal IV	Oryctolagus cuniculus	2.4.99.	AF121967	AAF28871.1	Q9N257	
OJ1217_F02.7	Oryza sativa (japonica cultivar- group)	n.d.	AP004084	BAD07616.1		_
OSJNBa0043L24.2 or OSJNBb0002J11.9	Oryza sativa (japonica cultivar- group)	n.d.	AL731626 AL662969	<b>CAD41185.1</b> CAE04714.1		
P0683f02.18 or P0489B03.1	Oryza sativa (japonica cultivar- group)	n.d.	AP003289 AP003794	<b>BAB63715.1</b> BAB90552.1		
α-2,6-sialyltransferase ST6GalNAc V (Siat7E) fragment)	Oryzias latipes	n.d.	AJ646876	CAG26705.1		
x-2,3-sialyltransferase ST3Gal I (Siat4)	Pan troglodytes	n.d.	AJ744803	CAG32839.1		
x-2,3-sialyltransferase 3T3Gal II (Siat5)	Pan troglodytes	n.d.	AJ744804	CAG32840.1		
x-2,3-sialyltransferase 5T3Gal III (Siat6)	Pan troglodytes	n.d.	AJ626819	CAF25177.1		
x-2,3-sialyltransferase 5T3Gal IV (Siat4c)	Pan troglodytes	n.d.	AJ626824	CAF25182.1		
x-2,3-sialyltransferase 3T3Gal VI (Siat10)	Pan troglodytes	n.d.	AJ744808	CAG32844.1		
x-2,6-sialyltransferase Sia7A)	Pan troglodytes	n.d.	AJ748740	CAG38615.1		_
x-2,6-sialyltransferase Sia7B)	Pan troglodytes	n.d.	AJ748741	CAG38616.1		
c-2,6-sialyltransferase T6GalNAc III (Siat7C)	Pan troglodytes	n.d.	AJ634454	CAG25676.1		
c-2,6-sialyltransferase T6GalNAc IV (Siat7D) ragment)	Pan troglodytes	n.d.	AJ646870	CAG26699.1		
r-2,6-sialyltransferase T6GalNAc V (Siat7E)	Pan troglodytes	n.d.	AJ646875	CAG26704.1		
c-2,6-sialyltransferase T6GalNAc VI (Siat7F) ragment)	Pan troglodytes	n.d.	AJ646882	CAG26711.1		
-2,8-sialyltransferase A (Siat8A)	Pan troglodytes	2.4.99.8	AJ697658	CAG26896.1		
-2,8-sialyltransferase 3 (Siat8B)	Pan troglodytes			CAG26897.1		
-2,8-sialyltransferase C (Siat8C)	Pan troglodytes	n.d.		CAG26898.1		
-2,8-sialyltransferase D (Siat8D)	Pan troglodytes		AJ697661	CAG26899.1		
-2,8-sialyltransferase	Pan troglodytes	n.d.	AJ697662	CAG26900.1		

Protein	Organism	EC#	GenBar	nk / GenPept	SwissProt	PDE / 3D
8E (Siat8E)						1.00
α-2,8-sialyltransferase 8F (Siat8F)	Pan troglodytes	n.d.	AJ697663	CAG26901.1		_
galactosamide α-2,6- sialyltransferase I (ST6Gal I; Siat1)	Pan troglodytes	2.4.99.1	AJ627624	CAF29492.1		
galactosamide α-2,6- sialyltransferase II (ST6Gal II)	Pan troglodytes	n.d.	AJ627625	CAF29493.1		
GM3 synthase ST3Gal V (Siat9)	Pan troglodytes	n.d.	AJ744807	CAG32843.1		
S138L	Rabbit fibroma virus Kasza	n.d.	NC_001266	NP_052025		
α-2,3-sialyltransferase ST3Gal III	Rattus norvegicus	2.4.99.6	M97754 NM_031697	AAA42146.1 7NP_113885.1	Q02734	
α-2,3-sialyltransferase ST3Gal IV (Siat4c)	Rattus norvegicus	n.d.	AJ626825	CAF25183.1		
α-2,3-sialyltransferase ST3Gal VI	Rattus norvegicus	n.d.	AJ626743	CAF25053.1		
α-2,6-sialyltransferase ST3Gal II	Rattus norvegicus	2.4.99	X76988 NM_031695	CAA54293.1 NP_113883.1	Q11205	
α-2,6-sialyltransferase ST6Gal I	Rattus norvegicus	2.4.99.1	M18769 M83143	AAA41196.1 AAB07233.1	P13721	-
α-2,6-sialyltransferase ST6GalNAc I (Siat7A)	Rattus norvegicus	n.d.	AJ634458	CAG25684.1		
α-2,6-sialyltransferase ST6GalNAc II (Siat7B)	Rattus norvegicus	n.d.	AJ634457	CAG25679.1		
α-2,6-sialyltransferase ST6GalNAc III	Rattus norvegicus			AAC42086.1 AAH72501.1 NP_061996.1	Q64686	
α-2,6-sialyltransferase ST6GalNAc IV (Siat7D) (fragment)	Rattus norvegicus	n.d.	AJ646871	CAG26700.1		
α-2,6-sialyltransferase ST6GalNAc V (Siat7E)	Rattus norvegicus	n.d.	AJ646872	CAG26701.1		
α-2,6-sialyltransferase ST6GalNAc VI (Siat7F) (fragment)	Rattus norvegicus	n.d.	AJ646881	CAG26710.1		
α-2,8-sialyltransferase (GD3 synthase) ST8Sia	Rattus norvegicus	2.4.99		AAC27541.1 BAA08213.1	P70554 P97713	
α-2,8-sialyltransferase (SIAT8E)	Rattus norvegicus	n.d.	AJ699422	CAG27884.1		$\neg$
α-2,8-sialyltransferase (SIAT8F)	Rattus norvegicus	n.d.	AJ699423	CAG27885.1		
α-2,8-sialyltransferase ST8Sia II	Rattus norvegicus	2.4.99 L		AAA42147.1 NP_476497.1	207977 264688	
α-2,8-sialyltransferase ST8Sia III	Rattus norvegicus 2	2.4.99 L		AAB50061.1 F NP_037161.1	97877	
x-2,8-sialyltransferase ST8Sia IV	Rattus norvegicus 2	2.4.99 L	J90215	AB49989.1 C	008563	
galactosamide α-2,6- ialyltransferase II ST6Gal II)	Rattus norvegicus r	n.d. A	J627626	CAF29494.1		
GM3 synthase ST3Gal	Rattus norvegicus n			BAA33492.1 C	88830	

# FIGURE 11J

•	FIGURE	113				
Protein	Organism	EC#	GenBa	ink / GenPept	SwissProt	PE / 3
sialyltransferase ST3Gal-I (Siat4A)	Rattus norvegicus	n.d.	AJ748840	CAG44449.1		1/3
α-2,3-sialyltransferase (St3Gal-II)	Silurana tropicalis	n.d.	AJ585763	CAE51387.1		
α-2,6-sialyltransferase (Siat7b)	Silurana tropicalis	n.d.	AJ620650	CAF05849.1		
α-2,6-sialyltransferase (St6galnac)	Strongylocentrotus purpuratus	n.d.	AJ699425	CAG27887.1		
α-2,3-sialyltransferase (ST3GAL-III)	Sus scrofa .	n.d.	AJ585765	CAE51389.1		
α-2,3-sialyltransferase (ST3GAL-IV)	Sus scrofa	n.d.	AJ584674	CAE48299.1		
α-2,3-sialyltransferase ST3Gal I	Sus scrofa	2.4.99.4	M97753	AAA31125.1	Q02745	
α-2,6-sialyltransferase (fragment) ST6Gal I	Sus scrofa	2.4.99.1	AF136746	AAD33059.1	Q9XSG8	
galactosamide α-2,6- sialyltransferase (ST6GalNAc-V)	Sus scrofa	n.d.	AJ620948	CAF06585.2		
sialyltransferase (fragment) ST6Gal I	sus scrofa	n.d.	AF041031	AAC15633.1	O62717	
ST6GALNAC-V	Sus scrofa	n.d.	AJ620948	CAF06585.1		
α-2,3-sialyltransferase (Siat5-r)	Takifugu rubripes	n.d.	AJ744805	CAG32841.1		
α-2,3-sialyltransferase ST3Gal I (Siat4)	Takifugu rubripes	n.d.	AJ626816	CAF25174.1		_
α-2,3-sialyltransferase ST3Gal II (Siat5) (fragment)	Takifugu rubripes	n.d.	AJ626817	CAF25175.1		
α-2,3-sialyltransferase ST3Gal III (Siat6)	Takifugu rubripes	n.d.	AJ626818	CAF25176.1		-
α-2,6-sialyltransferase ST6Gal I (Siat1)	Takifugu rubripes	n.d.	AJ744800	CAG32836.1		
α-2,6-sialyltransferase ST6GalNAc II (Siat7B)	Takifugu rubripes	n.d.	AJ634460	CAG25681.1		
α-2,6-sialyltransferase ST6GalNAc II B (Siat7B- elated)	Takifugu rubripes	n.d.	AJ634461	CAG25682.1		
x-2,6-sialyltransferase ST6GalNAc III (Siat7C) fragment)	Takifugu rubripes	n.d.	AJ634456	CAG25678.1		
x-2,6-sialyltransferase 3T6GalNAc IV (siat7D) fragment)	Takifugu rubripes	2.4.99.3	Y17466 AJ646869	CAB44338.1 CAG26698.1	29W6U6	
x-2,6-sialyltransferase T6GalNAc V (Siat7E) ragment)	Takifugu rubripes	n.d.	AJ646873	CAG26702.1		_
c-2,6-sialyltransferase T6GalNAc VI (Siat7F) ragment)	Takifugu rubripes	n.d.	AJ646880	CAG26709.1		
r-2,8-sialyltransferase T8Sia I (Siat 8A) ragment)	Takifugu rubripes	n.d.	AJ715534	CAG29373.1		
t-2,8-sialyltransferase T8Sia II (Siat 8B) ragment)	Takifugu rubripes	n.d.	AJ715538	CAG29377.1		
r-2,8-sialyltransferase T8Sia III (Siat 8C) ragment)	Takifugu rubripes	n.d.	AJ715541	CAG29380.1		
-2,8-sialyltransferase T8Sia IIIr (Siat 8Cr)		n.d.	\J715542	CAG29381.1		
-2,8-sialyltransferase F8Sia V (Siat 8E)	Takifugu rubripes	n.d.	J715547	CAG29386.1		$\exists$

### FIGURE 11K

Protein	Organism	EC#	GenBa	nk / GenPept	SwissProt	
(fragment)					<u> </u>	/ 3
α-2,8-sialyltransferase ST8Sia VI (Siat 8F) (fragment)	Takifugu rubripes	n.d.	AJ715549	CAG29388.1		
α-2,8-sialyltransferase ST8Sia VIr (Siat 8Fr)	Takifugu rubripes	n.d.	AJ715550	CAG29389.1		
α-2,3-sialyltransferase (Siat5-r)	Tetraodon nigroviridis	n.d.	AJ744806	CAG32842.1		
α-2,3-sialyltransferase ST3Gal I (Siat4)	Tetraodon nigroviridis	n.d.	AJ744802	CAG32838.1		
α-2,3-sialyltransferase ST3Gal III (Siat6)	Tetraodon nigroviridis	n.d.	AJ626822	CAF25180.1		
α-2,6-sialyltransferase ST6GalNAc II (Siat7B)	Tetraodon nigroviridis	n.d.	AJ634462	CAG25683.1		
κ-2,6-sialyltransferase ST6GalNAc V (Siat7E) (fragment)	Tetraodon nigroviridis	n.d.	AJ646879	CAG26708.1		
α-2,8-sialyltransferase ST8Sia I (Siat 8A) (fragment)	Tetraodon nigroviridis	n.d.	AJ715536	CAG29375.1		
α-2,8-sialyltransferase ST8Sia II (Siat 8B) (fragment)	Tetraodon nigroviridis	n.d.	AJ715537	CAG29376.1		
α-2,8-sialyltransferase ST8Sia III (Siat 8C) (fragment)	Tetraodon nigroviridis	n.d.	AJ715539	CAG29378.1		
α-2,8-sialyltransferase ST8Sia IIIr (Siat 8Cr) fragment)	Tetraodon nigroviridis	n.d.	AJ715540	CAG29379.1		-
α-2,8-sialyltransferase ST8Sia V (Siat 8E) fragment)	Tetraodon nigroviridis	n.d.	AJ715548	CAG29387.1		
α-2,3-sialyltransferase St3Gal-II)	Xenopus laevis	n.d.	AJ585762	CAE51386.1		
∞-2,3-sialyltransferase St3Gal-VI)	Xenopus laevis	n.d.	AJ585766	CAE51390.1		
x-2,3-sialyltransferase 6t3Gal-III (Siat6)	Xenopus laevis	n.d.	AJ585764 AJ626823	CAE51388.1 CAF25181.1		
x-2,8- olysialyltransferase x-2,8-sialyltransferase	Xenopus laevis				O93234	
T8Siα-I (Siat8A;GD3 ynthase)	Xenopus laevis	n.d.	AY272056 AY272057 AJ704562	AAQ16162.1 AAQ16163.1 CAG28695.1		
Jnknown (protein for IGC:81265)	Xenopus laevis	n.d.	BC068760	AAH68760.1		
x-2,3-sialyltransferase 3Gal-VI) x-2,3-sialyltransferase	Xenopus tropicalis	n.d.	AJ626744	CAF25054.1		
Siat4c) C-2,6-sialyltransferase	Xenopus tropicalis  Xenopus tropicalis		AJ622908	CAF22058.1		
T6GalNAc V (Siat7E)			AJ646878	CAG26707.1		
c-2,8-sialyltransferase T8Sia III (Siat 8C) ragment)	Xenopus tropicalis	n.d.	AJ715544	CAG29383.1		
galactosamide α-2,6- alyltransferase II ST6Gal II)	Xenopus tropicalis	n.d.	AJ627628	CAF29496.1		
ialytransferase St8Sial	Xenopus tropicalis	n.d.	AY652775	AAT67042		$\dashv$
oly-α-2,8-sialosyl alyltransferase (NeuS)	Escherichia coli K1	2.4 N	A76370	AAA24213.1 C	257269	$\dashv$
oly-α-2,8-sialosyl		2.4 N	76370 60598	AAA24213.1 CAA43053.1	2 <b>572</b> 2474	

### FIGURE 11K

Protein	Organism	EC#	GenBa	nk / GenPept	SwissPro	
α-2,8	Neisseria	2.4,-	M95053	AAA20478.1	Q51281	<u> </u>
polysialyltransferase SiaD	meningitidis B1940		X78068	CAA54985.1		
SynE	Neisseria meningitidis FAM18	n.d.	U75650	AAB53842.1	006435	
polysialyltransferase (SiaD)(fragment)	Neisseria meningitidis M1019	n.d.	AY234192	AAO85290.1		
SiaD (fragment)	Neisseria meningitidis M209	n.d.	AY281046	AAP34769.1		
SiaD (fragment)	Neisseria meningitidis M3045	n.d.	AY281044	AAP34767.1		
polysialyltransferase (SiaD)(fragment)	Neisseria meningitidis M3315	n.d.	AY234191	AAO85289.1		
SiaD (fragment)	Neisseria meningitidis M3515	n.d.	AY281047	AAP34770.1		
polysialyltransferase (SiaD)(fragment)	Neisseria meningitidis M4211	n.d.	AY234190	AAO85288.1		
SiaD (fragment)	Neisseria meningitidis M4642	n.d.	AY281048	AAP34771.1		
polysialyltransferase (SiaD)(fragment)	Neisseria meningitidis M5177	n.d.	AY234193	AAO85291.1		
SiaD	Neisseria meningitidis M5178	n.d.	AY281043	AAP34766.1		
SiaD (fragment)	Neisseria meningitidis M980	n.d.	AY281045	AAP34768.1		
NMB0067	Neisseria meningitidis MC58	n.d.	NC_003112	NP_273131		
Lst	Aeromonas punctata Sch3	n.d.	AF126256	AAS66624.1		
ORF2	Haemophilus influenzae A2	n.d.	M94855	AAA24979.1		
HI1699	Haemophilus influenzae Rd	n.d.	U32842 NC_000907	AAC23345.1 NP_439841.1		
∝-2,3-sialyltransferase	Neisseria gonorrhoeae F62	2.4.99.4	U60664	AAC44539.1 AAE67205.1	P72074	
α-2,3-sialyltransferase	Neisseria meningitidis 126E, NRCC 4010	2.4.99.4	U60662	AAC44544.2		-
α-2,3-sialyltransferase	Neisseria meningitidis 406Y, NRCC 4030	2.4.99.4	U60661	AAC44543.1		
α-2,3-sialyltransferase (NMB0922)	Neisseria meningitidis MC58	I .	AE002443	AAC44541.1 AAF41330.1 NP_273962.1	P72097	
NMA1118	Neisseria meningitidis Z2491		AL162755 NC 003116	CAB84380.1 NP 283887.1	Q9JUV5	
PM0508	Pasteurella multocida PM70	n.d.	AE006086	AAK02592.1 NP 245445.1	Q9CNC4	
WaaH	Salmonella enterica SARB25			AAM82550.1	Q8KS93	
WaaH		n.d.	AF519788	AAM82551.1	Q8KS92	
WaaH		n.d.	AF519789	AAM82552.1		$\dashv$
VaaH		n.d.	AF519790	AAM82553.1		$\dashv$
VaaH	· · · · · · · · · · · · · · · · · · ·	n.d.	AF519791	AAM82554.1	Q8KS91	$\dashv$
VaaH		n.d.	AF519793	AAM82556.1	Q8KS89	$\neg$
VaaH		n.d.	AF519792	AAM82555.1	Q8KS90	-+

### FIGURE 11L

WaaH   Salmonella enterica   N.d.   AF519779   AAM88840.1   Q8KS99   Salmonella enterica   N.d.   AF519781   AAM88840.1   Q8KS99   AAM88841.1   Q8KS96   AAM88841.1   Q8KS96   AAM88841.1   Q8KS96   AAM88841.1   Q8KS96   AAM88841.1   Q8KS96   AAM88841.1   Q8KS96   AAM88841.1   Q8KS99   AAM88881.1   Q8KS99	Protein	Organism	EC#	# GenBa	nk / GenPept	SwissProt Pl	
Waah   Salmonella enterica   S.ARC10V   Waah   (fragment)   Salmonella enterica   S.ARC13V   Salmonella enterica   S.ARC13V   Salmonella enterica   S.ARC13V   Salmonella enterica   S.ARC13V   Salmonella enterica   S.ARC13V   Salmonella enterica   S.ARC13V   Salmonella enterica   S.ARC13V   Salmonella enterica   S.ARC14V   Salmonella enterica   S.ARC15V   Salm		SARB8	┪	<del></del>	1		
SARC12   Salmonella enterica   Salmonella		Salmonella enterica SARC10V		AF519779	AAM88840.1	Q8KS99	
SARC.13  Salmonelle enterica   n.d.   AF519783   AAM88844.1   Q8KS97   SARC.14    Salmonelle enterica   n.d.   AF519784   AAM88844.1   Q8KS97   SARC.14    Salmonelle enterica   n.d.   AF519785   AAM88844.1   Q8KS95   SARC.15    Salmonella enterica   n.d.   AF519785   AAM88844.1   Q8KS95   SARC.15    Salmonella enterica   n.d.   AF519772   AAM88834.1   Q8KSA4   SARC.31    Salmonella enterica   n.d.   AF519773   AAM88834.1   Q8KSA4   SARC.31    Salmonella enterica   n.d.   AF519773   AAM88833.1   Q8KSA3   SARC.31    WaaH   Salmonella enterica   n.d.   AF519774   AAM88833.1   Q8KSA2   SARC.31    Salmonella enterica   n.d.   AF519775   AAM88837.1   Q8KSA2   SARC.31    Salmonella enterica   n.d.   AF519775   AAM88837.1   Q8KSA2   SARC.31    Salmonella enterica   n.d.   AF519777   AAM88838.1   Q8KSA1   SARC.31    Salmonella enterica   n.d.   AF519778   AAM88838.1   Q8KSA1   SARC.31    Salmonella enterica   n.d.   AF519778   AAM88839.1   Q8KSA2   SARC.32    Salmonella enterica   n.d.   AF519778   AAM88839.1   Q8KSA2   SARC.32    Salmonella enterica   n.d.   AF519778   AAM88839.1   Q8KSA2   SARC.32    Salmonella enterica   n.d.   AF519778   AAM88839.1   Q8KSA2   SARC.32    Salmonella enterica   n.d.   AF519778   AAM88839.1   Q8KSA2   SARC.32    Salmonella enterica   n.d.   AF519778   AAM88839.1   Q8KSA2   SARC.32    Salmonella enterica   n.d.   AF401529   AAL06004.1   Q93C25   Salmonella enterica   n.d.   AF401529   AAL06004.1   Q93C25   Salmonella enterica   n.d.   AF401529   AAL06004.1   Q93C25   Salmonella enterica   n.d.   AF400048   AAK73183.1   Salmonella enterica   n.d.   AF400048   AAK73183.1   Salmonella enterica   Salmonella enterica   n.d.   AF400048   AAK73183.1   Salmonella enterica   Salmonella enterica   n.d.   AF400048   AAK73183.1   Salmonella enterica   Salmonella enterica   n.d.   AF400048   AAK73183.1   Salmonella enterica   Salmonella enterica   n.d.   AF400048   AAK73183.1   Salmonella enterica   Salmonella enterica   N.d.   AF400048   AAK73183.1   Salmonella enterica   Salmonella enterica   N.			n.d.	AF519781	AAM88842.1		
WaaH (fragment)   Salmonella enterica   n.d.   AF519783   AAM88844.1   Q8KS97   SARC141   Salmonella enterica   n.d.   AF519784   AAM88845.1   Q8KS95   ARC1511   Salmonella enterica   n.d.   AF519785   AAM88846.1   Q8KS95   ARC1511   Salmonella enterica   n.d.   AF519772   AAM88834.1   Q8KSA5   ARC1511   Salmonella enterica   n.d.   AF519772   AAM88834.1   Q8KSA5   ARC313   Salmonella enterica   n.d.   AF519773   AAM88835.1   Q8KSA5   ARC314   Salmonella enterica   n.d.   AF519773   AAM88835.1   Q8KSA5   ARC5111   ARC5111   ARM88835.1   Q8KSA5   ARC5111   ARM88835.1   Q8KSA5   ARC5111   ARM88835.1   Q8KSA5   ARC5111   ARM88835.1   Q8KSA5   ARC6111   ARM88835.1   Q8KSA5   ARC6111   ARM88835.1   Q8KSA5   ARC6111   ARM88835.1   Q8KSA5   ARC6111   ARM88835.1   Q8KSA5   ARC6111   ARM88835.1   Q8KSA5   ARC6111   ARM88835.1   Q8KSA5   ARC6111   ARM88835.1   Q8KSA5   ARC6111   ARM88835.1   Q8KSA5   ARC6111   ARM88835.1   Q8KSA5   ARC6111   ARM88835.1   Q8KSA5   ARC6111   ARM88835.1   Q8KSA5   ARC6111   ARM88835.1   Q8KSA5   ARC6111   ARM88835.1   Q8KSA5   ARC6111   ARM88835.1   Q8KSA5   ARC6111   ARM88835.1   Q8KSA5   ARC6111   ARC6111   ARM88835.1   Q8KSA5   ARC6111   ARM88835.1   Q8KSA5   ARC6111   ARM88835.1   Q8KSA5   ARC6111   ARM88835.1   Q8KSA5   ARC6111   ARM88835.1   Q8KSA5   ARC6111   ARM88835.1   Q8KSA5   ARC6111   ARM88835.1   Q8KSA5   ARC6111   ARC6111   ARC6111   ARC6111   ARM88835.1   Q8KSA5   ARC6111   ARC6111   ARC6111   ARC6111   ARC6111   ARC6111   ARC6111   ARC6111   ARC6111   ARC6111   ARM88835.1   Q8KSA5   ARC6111   ARC6111   ARC6111   ARM88835.1   Q8KSA5   ARM88835.1   Q8KSA5   ARC611   ARM88835.1   Q8KSA5   ARM88835.1   Q8KSA5   ARC611   ARM88835.1   Q8KSA5   ARM88835.1   Q8KSA5   ARM88835.1   Q8KSA5   ARC6111   ARC6111   ARC6111   ARC6111   ARM88835.1   Q8KSA5   ARC6111   ARC			n.d.	AF519782	AAM88843.1	Q8KS98	
WaaH   Salmonella enterica   S.ARC.15			n.d.	AF519783	AAM88844.1	Q8KS97	
WaaH (fragment)   Salmonella enterica   S	WaaH		n.d.	AF519784	AAM88845.1	Q8KS96	
SARC3  Salmonella enterica   n.d.   AF519773   AAM88835.1   Q8KSA3   SARC4    Salmonella enterica   n.d.   AF519774   AAM88835.1   Q8KSA3   SARC5    SARC5    SARC5    SARC5    SARC5    SARC5    SARC5    SARC5    SARC6    SARC			n.d.	AF519785	AAM88846.1	Q8KS95	
SARC4  Salmonella enterica   Salmonella   Sal			n.d.	AF519772	AAM88834.1	Q8KSA4	
SARC5Ila   Salmonella enterica   Salmonel	WaaH (fragment)		n.d.	AF519773	AAM88835.1	Q8KSA3	
WaaH	WaaH	Salmonella enterica	n.d.	AF519774	AAM88836.1		
WaaH   Salmonella enterica   n.d.   AF519777   AAM88838.1   Q8KSA1   WaaH   Salmonella enterica   SARC9   N.d.   AF519778   AAM88839.1   Q8KSA0   UDP-glucose : α-1.2-   glucosyltransferase (WaaH)   Salmonella enterica   SARC9   Salmonella enterica   SARC9   Salmonella enterica   SARC   Salmonella enterica   SARC   Salmonella enterica   SARC   Salmonella enterica   SARC   Salmonella enterica   SARC   Salmonella enterica   SARC   Salmonella enterica   SARC   Salmonella enterica   SARC   Salmonella enterica   SARC   Salmonella enterica   SARC   Salmonella enterica   SARC   Salmonella enterica   Salmonella enterica   SARC   Salmonella enterica   Salmone	WaaH	Salmonella enterica	n.d.	AF519775	AAM88837.1	Q8KSA2	
SARC9V   Salmonella enterica subsp. arizonae (WaaH)   Salmonella enterica subsp. arizonae (WaaH)   Salmonella enterica subsp. arizonae (SARC 5   Salmonella en	WaaH	Salmonella enterica	n.d.	AF519777	AAM88838.1	Q8KSA1	
UDP-glucose : α-1,2-glucose	WaaH	Salmonella enterica	n.d.	AF519778	AAM88839.1	Q8KSA0	
	glucosyltransferase (WaaH)	Salmonella enterica subsp. arizonae	2.4.1	AF511116	AAM48166.1		
Campylobacter   igiuni 81-176   Campylobacter   igiuni 81-176   Campylobacter   igiuni 81-176   Campylobacter   igiuni ATCC 43429   Campylobacter   igiuni ATCC 43430   Campylobacter   igiuni ATCC 43430   Campylobacter   igiuni ATCC 43430   Campylobacter   igiuni ATCC 43430   Campylobacter   igiuni ATCC 43430   Campylobacter   igiuni ATCC 43430   Campylobacter   igiuni ATCC 43438   Campylobacter   igiuni ATCC 43446   Campylobacter   igiuni ATCC 43446   Campylobacter   igiuni ATCC 43446   Campylobacter   igiuni ATCC 43456   Campylobacter   igiuni ATCC 43456   Campylobacter   igiuni ATCC 43456   Campylobacter   igiuni ATCC 43456   Campylobacter   igiuni ATCC 43456   Campylobacter   igiuni ATCC 43456   Campylobacter   igiuni ATCC 43460   Campylobacter   igiuni ATCC 43460   Campylobacter   igiuni ATCC 43460   Campylobacter   igiuni ATCC 43460   Campylobacter   igiuni ATCC 43460   Campylobacter   igiuni ATCC 43460   Campylobacter   igiuni ATCC 43460   Campylobacter   igiuni ATCC 43460   Campylobacter   igiuni ATCC 700297   AAR36462.1   igiuni ATCC 700297   Campylobacter   igiuni ATCC 700297   Campylobacter   igiuni ATCC 700297   Campylobacter   igiuni ATCC 700297   Campylobacter   igiuni MSC57360   Campylobacter   igiuni MSC57360   Campylobacter   igiuni NCTC 11168   Campylobacter   igiuni NCTC 11168   Campylobacter   igiuni NCTC 11168   Campylobacter   igiuni NCTC 11168   Campylobacter   igiuni NCTC 11168   Campylobacter   igiuni NCTC 11168   Campylobacter   igiuni O:10   AX934427   CAF04169.1   CAF0416			n.d.	AF401529	AAL06004.1	Q93CZ5	
α-2,3-sialyltransferase (Cst-III)         Campylobacter jejuni ATCC 43429         2.4.99         AY044156         AAK73183.1           α-2,3-sialyltransferase (Cst-III)         Campylobacter jejuni ATCC 43430         2.4.99         AF400047         AAK85419.1           α-2,3-sialyltransferase (Cst-III)         Campylobacter jejuni ATCC 43432         2.4.99         AF215659         AAG43979.1         Q9F0M9           α-2,3/8- sialyltransferase (St-III)         Campylobacter jejuni ATCC 43446         1.0.         AF400048         AAK91725.1         Q93MQ0           α-2,3-sialyltransferase (St-III)         Campylobacter jejuni ATCC 43446         2.4.99         AF167344         AAF34137.1	Cst	Campylobacter	n.d.	AF305571	AAL09368.1		
	Cst-III)	Campylobacter jejuni ATCC 43429	2.4.99	AY044156	AAK73183.1		
α-2,3-sialyltransferase (Cst-II)         Campylobacter igiuni ATCC 43432         2.4.99         AF215659         AAG43979.1         Q9F0M9           α-2,3/β- sialyltransferase (CstII)         Campylobacter igiuni ATCC 43438         n.d.         AF400048         AAK91725.1         Q93MQ0           α-2,3-sialyltransferase cst-II         Campylobacter igiuni ATCC 43446         2.4.99         AF167344         AAF34137.1         Q93D05           α-2,3-α-2,8- isalyltransferase (CstII)         Campylobacter igiuni ATCC 43460         2.4.99         AF401528         AAL05990.1         Q93D05           α-2,3/α-2,β- isalyltransferase (Cst-II)         Campylobacter igiuni ATCC 700297         n.d.         AF216647         AAL36462.1           α-2,3-sialyltransferase (Cst-III)         Campylobacter igiuni MSC57360         n.d.         AY422197         AAR82875.1           α-2,3-sialyltransferase still (j1140         Campylobacter igiuni NCTC 11168         2.4.99         AF195055         AAG29922.1           α-2,3/α-2,β- ialyltransferase II (cstII)         Campylobacter igiuni O:10         n.d.         -         -         AAO96669.1           α-2,3/α-2,β- ialyltransferase II (cstII)         Campylobacter igiuni O:19         n.d.         AX934431         CAF04169.1	Cst-III)	jejuni ATCC 43430	2.4.99	AF400047	AAK85419.1		
igiuni ATCC 43438   Campylobacter   igiuni ATCC 43446   Campylobacter   igiuni ATCC 43446   Campylobacter   igiuni ATCC 43446   Campylobacter   igiuni ATCC 43446   Campylobacter   igiuni ATCC 43446   Campylobacter   igiuni ATCC 43456   Campylobacter   igiuni ATCC 43456   Campylobacter   igiuni ATCC 43456   Campylobacter   igiuni ATCC 43460   Campylobacter   igiuni ATCC 43460   Campylobacter   igiuni ATCC 43460   Campylobacter   igiuni ATCC 700297   Campylobacter   igiuni ATCC 700297   Campylobacter   igiuni ATCC 700297   Campylobacter   igiuni ATCC 700297   Campylobacter   igiuni ATCC 700297   Campylobacter   igiuni MSC57360   Campylobacter   igiuni MSC57360   Campylobacter   igiuni MSC57360   Campylobacter   igiuni NCTC 11168   Campylobacter   igiuni NCTC 11168   Campylobacter   igiuni NCTC 11168   Campylobacter   igiuni NCTC 11168   Campylobacter   igiuni O:10   Campylobacter   igiuni O:10   Campylobacter   igiuni O:19   Campylobacter   igiuni O:10   Campylobacter   igiuni O:10   Campylobacter   igiuni O:1	Cst-II)	Campylobacter	2.4.99	AF215659	AAG43979.1	Q9F0M9	
α-2,3-sialyltransferase cst-II         Campylobacter jejuni ATCC 43446         2.4.99 AF167344         AAF34137.1           α-2,3-sialyltransferase Cst-II)         Campylobacter jejuni ATCC 43456         2.4.99 AF401528         AAL05990.1         Q93D05           α-2,3-α-2,8-sialyltransferase (CstII)         Campylobacter jejuni ATCC 43460         2.4.99 AF216647         AAL36462.1         Q938X6           α-2,3/8-sialyltransferase (Cst-III)         Campylobacter jejuni ATCC 700297         n.d. AF216647         AAR82875.1         AAR82875.1           α-2,3-sialyltransferase stIII         Campylobacter jejuni MSC57360         2.4.99 AF195055         AAG29922.1           α-2,3-sialyltransferase stIII cj1140         Campylobacter jejuni NCTC 11168         2.4.99 AL139077         CAB73395.1 NP 282288.1         Q9PNF4           α-2,3/α-2,8- ialyltransferase II (cstIII)         Campylobacter jejuni O:10         AAO96669.1         AAO96669.1           α-2,3/α-2,8- ialyltransferase II (cstIII)         Campylobacter jejuni O:19         n.d. AX934431         CAF04169.1			n.d.	AF400048	AAK91725.1	Q93MQ0	
igjuni ATCC 43456   Campylobacter   igjuni ATCC 43456   Campylobacter   igjuni ATCC 43460   Campylobacter   igjuni ATCC 43460   Campylobacter   igjuni ATCC 43460   Campylobacter   igjuni ATCC 700297   Campylobacter   igjuni ATCC 700297   Campylobacter   igjuni GB11   Campylobacter   igjuni MSC57360   Campylobacter   igjuni MSC57360   Campylobacter   igjuni NCTC 11168   Campylobacter   igjuni NCTC 11168   Campylobacter   igjuni NCTC 11168   Campylobacter   igjuni O:10   Campylobacter   igjuni O:10   Campylobacter   igjuni O:19   Campylobacter   igjuni O:10   Campylobacter   igjuni O:10   Campylobacter   igjuni O:10   Campylobacter   igjuni O:10   Campylobacter   igjuni O:10   Campylobacter   igjuni O:10   Campylobacter   igjuni O:10   Campylobacter   igjuni O:10   Campylobacter   igjuni O:10   Campylobacter   igjuni O:10   Campylobacter   igjuni O:10   Campylobacter   igjuni O:10   Campylobacter   igjuni O:10   Campylobacter   igjuni O:10   Campylobacter   igjuni O:10   Campylobacter   igjuni O:10   Campyloba		Campylobacter	2.4.99	AF167344	AAF34137.1		
α-2,3-/α-2,8-sialyltransferase (Cstill)         Campylobacter jejuni ATCC 43460         2.4.99 AY044868         AAK96001.1         Q938X6           α-2,3/8-sialyltransferase (Cst-II)         Campylobacter jejuni ATCC 700297         n.d. AF216647         AAL36462.1           ORF         Campylobacter jejuni GB11         n.d. AY422197         AAR82875.1           α-2,3-sialyltransferase stIII         Campylobacter jejuni MSC57360         2.4.99 AF195055         AAG29922.1           α-2,3-sialyltransferase stIII Cj1140         Campylobacter jejuni NCTC 11168         2.4.99 AL139077         CAB73395.1 Q9PNF4           α-2,3/α-2,8- ialyltransferase II (cstII)         Campylobacter jejuni O:10         n.d. AAO96669.1         AAO96669.1           α-2,3/α-2,8- ialyltransferase II (cstII)         Campylobacter jejuni O:19         n.d. AX934431         CAF04169.1	Cst-II)	Campylobacter	2.4.99	AF401528	AAL05990.1	Q93D05	
Campylobacter   jejuni ATCC 700297   n.d.   AF216647   AAL36462.1     Campylobacter   jejuni GB11   n.d.   AY422197   AAR82875.1     Campylobacter   jejuni GB11   2.4.99   AF195055   AAG29922.1     Campylobacter   jejuni MSC57360   2.4.99   AL139077   CAB73395.1   Q9PNF4     Still Cj1140   Campylobacter   jejuni NCTC 11168   NC 002163   NP 282288.1     Campylobacter   jejuni O:10   AX934427   CAF04167.1     Campylobacter   jejuni O:19   n.d.   AX934431   CAF04169.1     Campylobacter   jejuni O:19   N.d.   AX934431   CAF04169.1     Campylobacter   jejuni O:19   N.d.   AX934431   CAF04169.1     Campylobacter   jejuni O:19   N.d.   AX934431   CAF04169.1     Campylobacter   jejuni O:19   N.d.   AX934431   CAF04169.1     Campylobacter   jejuni O:19   N.d.   AX934431   CAF04169.1     Campylobacter   jejuni O:19   N.d.   AX934431   CAF04169.1     Campylobacter   jejuni O:19   N.d.   AX934431   CAF04169.1     Campylobacter   jejuni O:19   N.d.   AX934431   CAF04169.1     Campylobacter   jejuni O:19   N.d.   AX934431   CAF04169.1     Campylobacter   jejuni O:19   N.d.   AX934431   CAF04169.1     Campylobacter   jejuni O:19   N.d.   AX934431   CAF04169.1     Campylobacter   jejuni O:19   N.d.   AX934431   CAF04169.1     Campylobacter   jejuni O:19   N.d.   AX934431   CAF04169.1     Campylobacter   jejuni O:19   N.d.   AX934431   CAF04169.1     Campylobacter   jejuni O:19   N.d.   AX934431   CAF04169.1     Campylobacter   jejuni O:19   N.d.   AX934431   CAF04169.1     Campylobacter   jejuni O:19   N.d.   AX934431   CAF04169.1     Campylobacter   jejuni O:19   N.d.   AX934431   N.d.   AX934431   N.d.   AX934431   N.d.   AX934431     Campylobacter   Jejuni O:19   N.d.   AX934431		Campylobacter	2.4.99	AY044868	AAK96001.1	Q938X6	
Campylobacter   jejuni GB11   Campylobacter   jejuni GB11   Campylobacter   jejuni MSC57360   Campylobacter   jejuni MSC57360   Campylobacter   jejuni MSC57360   Campylobacter   jejuni NCTC 11168   Campylobacter   jejuni NCTC 11168   Campylobacter   jejuni NCTC 11168   Campylobacter   jejuni O:10   CAF04167.1   CAF04167.1   CAF04169.1   jejuni O:19   CAF04169.1   Jejuni O:19   CAF04169.1   CAF04169.1   Jejuni O:19   CAF04169.1   CAF04169.1   Jejuni O:19   CAF04169.1   CAF04169.1   CAF04169.1   Jejuni O:19   CAF04169.1   CAF04169.		Campylobacter	n.d.	AF216647	AAL36462.1		
Campylobacter   jejuni MSC57360   CAB73395.1   Q9PNF4   StIII   Cj1140   Campylobacter   jejuni NCTC 11168   Campylobacter   jejuni NCTC 11168   Campylobacter   jejuni NCTC 11168   Campylobacter   Jejuni O:10   CAB73395.1   Q9PNF4   NC 002163   NP 282288.1   Campylobacter   Jejuni O:10   CAF04167.1   CAF04167.1   CAF04169.1   Jejuni O:19   CAF04169.1   Jejuni O:19   CAF04169.1		Campylobacter	n.d.	AY422197	AAR82875.1		
Campylobacter   jejuni NCTC 11168   Campylobacter   jejuni NCTC 11168   Campylobacter   NC 002163   NP 282288.1     x-2,3/α-2,8-   ialyltransferase   (cstii)   (campylobacter   jejuni O:10   Campylobacter   jejuni O:10   Campylobacter   jejuni O:19   Campylobacter   jejuni O:19   Campylobacter   jejuni O:19   Campylobacter   jejuni O:19   Campylobacter   jejuni O:19   Campylobacter   jejuni O:19   Campylobacter   jejuni O:19   Campylobacter   jejuni O:19   Campylobacter   jejuni O:19   Campylobacter   jejuni O:19   Campylobacter   jejuni O:19   Campylobacter   jejuni O:19   Campylobacter   jejuni O:19   Campylobacter   jejuni O:19   Campylobacter   jejuni O:19   Campylobacter   jejuni O:19   Campylobacter   jejuni O:10   Campylobacter   jejuni		Campylobacter	2.4.99	AF195055	AAG29922.1		
x-2,3/α-2,8- ialyltransferase II (cstII)  x-2,3/α-2,8- ialyltransferase II (cstII)  x-2,3/α-2,8- ialyltransferase II  Campylobacter jejuni O:19  CAF04169.1  CAF04169.1  CStII)		Campylobacter		Y .		9PNF4	
x-2,3/α-2,8- ialyltransferase II CstII)  Campylobacter i.d. AX934431 CAF04169.1 jejuni O:19	c-2,3/α-2,8-	Campylobacter r	n.d.	-	AAO96669.1		
7.2.2 m 2.9	c-2,3/α-2,8- alyltransferase II OstII)	Campylobacter r					
Campylobacter n.d. AX934436 CAF04171.1 jejuni O:36 CAF04170.1 CAF04170.1	t-2,3/α-2,8- alyltransferase II CstII)	jejuni O:36	n.d.	AX934436	CAF04171.1		

### FIGURE 11M

Protein	Organism	EC#	GenBank / GenPept		SwissPro	ot PDE
sialyltransferase II (CstII)	jejuni 0:4					7 30
α-2,3/α-2,8- sialyltransferase II (CstII)	Campylobacter jejuni O:41	n.d.	- - AX934429	AAO96670.1 AAT17967.1 CAF04168.1		
α-2,3-sialyltransferase	Campylobacter jejuni OH4384	2.4.99	AF130466		Q9RGF1	
bifunctional α-2,3/-2,8- sialyltransferase (Cst-II)	Campylobacter jejuni OH4384	2.4.99	AF130984 AX934425	AAF31771.1	1RO7 1RO8	C A
HI0352 (fragment)	Haemophilus influenzae Rd	n.d.	U32720 X57315 NC 000907	CAA40567.1	P24324	•
PM1174	Pasteurella multocida PM70	n.d.	AE006157 NC_002663	AAK03258.1	Q9CLP3	
Sequence 10 from patent US 6503744	Unknown.	n.d.		AAO96672.1		
Sequence 10 from patent US 6699705	Unknown.	n.d.	_	AAT17969.1		
Sequence 12 from patent US 6699705	Unknown.	n.d.	-	AAT17970.1		
Sequence 2 from patent US 6709834	Unknown.	n.d.	-	AAT23232.1		
Sequence 3 from patent US 6503744	Unknown.	n.d.	•	AAO96668.1		
Sequence 3 from patent JS 6699705	Unknown.	n.d.	-	AAT17965.1		
Sequence 34 from patent US 6503744	Unknown.	n.d.	-	AAO96684.1		
Sequence 35 from patent US 6503744 fragment)	Unknown.	n.d.		AAO96685.1 AAS36262.1		
Sequence 48 from patent US 6699705	Unknown.	n.d		AAT17988.1		
Sequence 5 from patent IS 6699705	Unknown.	n.d.		AAT17966.1		
Sequence 9 from patent JS 6503744	Unknown.	n.d		AAO96671.1	·	